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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
10/773,654	02/05/2004	George N. Cox III	4152-1-PUS-6	7639
22442 75	590 08/03/2006		EXAMINER	
SHERIDAN ROSS PC			STOICA, ELLY GERALD	
1560 BROADWAY SUITE 1200		ART UNIT	PAPER NUMBER	
DENVER, CO	80202		1647	
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Please find below and/or attached an Office communication concerning this application or proceeding.

Paper No(s)/Mail Date _____.

6) Other: ____.

Detailed Action

Information Disclosure Statement

1. The information disclosure statement (IDS) submitted on February 5, 2004 is in compliance with the provisions of 37 CFR 1.97. Accordingly, the information disclosure statement is being considered by the examiner.

Priority

2. Applicant's claim for the benefit of to a prior application 10/400377 is acknowledged.

Status of the claims

3. Currently, claims 24-46 are pending in the application. In the preliminary amendment filed on February 5, 2004, the claims 1-23 were cancelled by the applicant.

Claim Rejections - 35 USC § 112

4. The following is a quotation of the first paragraph of 35 U.S.C. 112:

The specification shall contain a written description of the invention, and of the manner and process of making and using it, in such full, clear, concise, and exact terms as to enable any person skilled in the art to which it pertains, or with which it is most nearly connected, to make and use the same and shall set forth the best mode contemplated by the inventor of carrying out his invention.

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5. Claims 24-46 rejected under 35 U.S.C. 112, first paragraph, as failing to comply with the enablement requirement. The claims contain subject matter which was not described in the specification in such a way as to enable one skilled in the art to which it pertains, or with which it is most nearly connected, to make and/or use the invention. The claims are drawn to a genus of granulocyte-macrophage colony-stimulating factor (GM-CSF) variants described as having a cysteine residue inserted before, after or between certain amino acid residues and having an in vitro ability to induce GM-CSF induced cell proliferation.

The factors considered when determining if the disclosure satisfies the enablement requirement and whether any necessary experimentation is "undue" include, but are not limited to:

1) nature of the invention, 2) state of the prior art, 3) relative skill of those in the art, 4) level of predictability in the art, 5) existence of working examples, 6) breadth of claims, 7) amount of direction or guidance by the inventor, and 8) quantity of experimentation needed to make or use the invention. In re Wands, 858 F.2d 731, 737, 8 USPQ2d 1400, 1404 (Fed. Cir. 1988).

In the present case, the Applicant has provided a structural limitation (insertion of a cysteine before, after or between certain amino acid residues of GM-CSF) and a functional limitation (the cell proliferation requirement). The cysteines are needed for PEGylation of the modified growth factor for increase of the half-life of the insertion variants of the GM-CSF claimed. The state of the prior art is adequate for the conceptual construction of cysteine mutants or PEGylated growth factors (Cunningham

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and Wells, Science 244, 1081-1085, Goodson and Katre, Biotechnology 1990, 343-6). Even though the relative skill existent in the art at the time of the priority date claimed would permit the making of certain cysteine mutants for Growth Hormone (US Pat. 6,608183), in the specification no data are presented as to mutants formed by insertion of cysteine residues in the structure of the GM-CSF. There are teachings in the art that underscore the uncertainty in protein modification in general and in the effects of modifying any particular residue in a protein sequence absent specific teachings relating the amino acid to the protein's function and structure (Bowie et al., Science 247, 1306-131). In the specification, the Applicant proposes "rules" for the modification of the target proteins, members of the Growth Hormone super family, (pages 11 and 12 of the specification) with three key components.

First, the specification identifies as preferred sites for modification those regions of the Growth Hormone supergene family corresponding to the pre-Helix A region, and the region distal to the last helix of the protein, and the A-B, the B-C, the C-D loops (i.e. the loop regions) of the proteins (page 10, lines 10-28 and page 11, lines 14-17)

Second, the application additionally indicates that N- and 0- glycosylation sites may also be preferred sites for protein modification (page 11, lines 19-22). Finally, the application teaches that these rules may be applied to other proteins, and that in such other proteins the amino acids that can be replaced with cysteine without significant loss of "biological activity", as is the insertion of cysteine between two amino acids that are situated in the "disclosed" regions (p11, lines 18-19). It is also noted that the application and the claims include as potential modification sites the first three and the last three

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residues in each of the A, B, C, and D helices (page 3, lines 3-20 and claim 26), although the application does not refer to these regions in the description of the "rules" on pages 11 and 12.

It is noted that, "where there is unpredictability in performance of certain species or subcombinations other than those specifically enumerated, one skilled in the art may be found not to have been placed in possession of a genus or combination claimed" In re Smyth, 178 U.S.P.Q. 279 at 284-85 (CCPA 1973). Thus, where there is uncertainty in the art, even the presence of multiple species within a claimed genus does not necessarily demonstrate possession of the genus. In the present case, the state of the art has provided a good deal of evidence supporting a finding of uncertainty in the art. However, the application provides only the teachings of the indicated "rules" to support the present claims to the genus. The rules and disclosed species are not deemed sufficient to overcome the uncertainties in the art. Several teaches in the art demonstrate that:

-modification of certain amino acids found within the pre-alpha helix region of the protein results in loss of protein activity;

-amino acid deletion is not predictive of the effects of substitution modifications to the same residues;

-insertion of cysteine has a greater likelihood of disrupting protein structure.

Thus, significant evidence of uncertainty has been presented. The embodiment described in the specification and claimed is rather a prophetic one based on predicted results rather than work actually conducted or results actually achieved.

The strongest contradiction between the teachings in the art and the applicant proposed "rules" are the teachings of Shaw (U.S. 5,166,322) and Zurawski et al (EMBO Journal 12: 5113-5119). The last reference demonstrates that many "unimportant residues" in the GH superfamily protein IL-2 were intolerant to cysteine modifications. It is noted that an alignment of the IL-2 reference in Figure 1 of Zurawski with the structural teachings of IL-2 found in Bazan et al., (Science 257: 410-13) indicates that the A, B, C, and D helices of the portion of IL-2 correspond, respectively, to the following residues of the Figure 1 sequence: A, R41 (only the C-terminal residue of this helix shown); B, K.68-187; C, N99-K1 12; and D, V130-5142 (all of D helix not shown). Taking into consideration the teachings of the Bazan reference, the Zurawski reference indicates that certain residues found in the A-B loop, the last three residues of each of the A and C helices, and in the first three residues of the C helix are found among those residues described as intolerant to cysteine substitutions. Thus, the reference supports the assertion that those in the art would not be able to predict, based on the teachings in the prior art, which amino residues would be tolerant to cysteine modification according to the teachings of Braxton (US patent 5,766,897). However, these same teachings also demonstrate that cysteine substitutions may not be made freely among the residues in various regions identified by the first "rule" presented in the application. In view of these teachings, it appears that the application of the Applicant's first rule would still result in uncertainty in the operability of any particular GM-CSF variant.

The second "rule" provided by the application is similarly contradicted by the teachings

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of the Shaw reference. This reference "found that the glycosylation site in IL-3 (asparagine 15) is not useful for creating cysteine mutants in IL-3 because the mutant protein is not biologically active." IL-3 is also among the proteins identified in the application as a member of the GH superfamily. Thus, the teachings of this reference also demonstrate uncertainty in operability of protein variants even upon the application of the second rule. Moreover, the art, and in particular Zurawski, indicate that the effect of the cysteine substations varies with amino acid being substituted. For example, residue 42 (in the A-B loop) of IL-2 was indicated to be intolerant to cysteine substitution, whereas the substitution of amino acids 43 and 44 had relatively little effect on the protein activity. While the claims include both structural and functional requirements, the teachings of the Zurawski reference indicate that the structural requirements fail to correlate with the functional requirements. In view of the lack of correlation between the structures and functions relied on to describe the claimed genus, and the evidence of uncertainty in the operability of any particular species of the claimed invention, even upon the application of the "rules", the teachings of the application are not deemed sufficient to provide descriptive support for the claimed genus of GM-CSF variants, without undue experimentation needed to obtain the particular bio-active GM-CSF insertion mutants.

A conclusion of lack of enablement means that, based on the evidence regarding each of the above factors, the specification, at the time the application was filed, would not have taught one skilled in the art how to make and/or use the full scope of the claimed

invention without undue experimentation. In re Wright, 999 F.2d 1557,1562, 27 USPQ2d 1510, 1513 (Fed. Cir. 1993).

The Applicant is encouraged to provide any evidence to demonstrate that the disclosure enables the claimed invention.

Conclusion

- 6. No claims are allowed.
- 7. Any inquiry concerning this communication or earlier communications from the examiner should be directed to Elly-Gerald Stoica whose telephone number is (571) 272-9941. The examiner can normally be reached on 8:30-17:00.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Brenda G. Brumback can be reached on (571) 272-0961. The fax phone number for the organization where this application or proceeding is assigned is 571-273-8300.

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Business Center (EBC) at 866-217-9197 (toll-free). If you would like assistance from a USPTO Customer Service Representative or access to the automated information system, call 800-786-9199 (IN USA OR CANADA) or 571-272-1000.

LORRAINE SPECTOR PRIMARY EXAMINER